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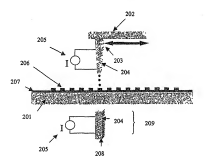
(71) Demandeur/Applicant: NANO-PROPRIETARY, INC., US

(72) Inventeurs/Inventors:

MAO, DONGSHENG, US; YANIV, ZVI, US; FINK, RICHARD LEE, US

(74) Agent: KIRBY EADES GALE BAKER

(54) Titre: METALLISATION DE NANOTUBES DE CARBONE POUR DES APPLICATIONS D'EMISSION DE CHAMP (54) Title: METALLIZATION OF CARBON NANOTUBES FOR FIELD EMISSION APPLICATIONS



(57) Abrégé/Abstract:

The present invention is directed towards metalized carbon nanotubes, methods for making metalized carbon nanotubes using an electroless plating technique, methods for disprising metalized carbon nanotubes onto a substrate, and methods for aligning magnetically-active metalized carbon nanotubes. The present invention is also directed towards cold carbode field emitting materials comprising metalized carbon nanotubes, and methods of using metalized carbon nanotubes as cold carbode field emitting.



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(71) Applicant: SI DIAMOND TECHNOLOGY, RNC.

[USJUS]; 3006, Longhorn, Blvd., Suite 107, Austin, TX 78758 (US).
 [72] Inventors: MAO, DONGSHENG; 8617 Spicowood Springs Road, 423, Austin, TX 78759 (US), YANNY, Zvi., 5810 Long Court, Austin, TX 78730 (US). FINNS, Richard, Los 3909 Hawkshead Drive, Austin, TX 78737

(74) Agents: KORDZIK, Kelly, K. et al.; WINSTBAD SECHREST & MINICK P.C., P.O.Box 50784, 1201 Main Street, Dallas, TX 75250-0784 (US).

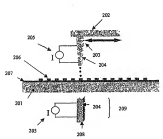
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## (54) Title: METALLIZATION OF CARBON NANOTUBES FOR FIELD EMISSION APPLICATIONS



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\*\*The present invention is a method for a single metallized cutton associates are cold cathods field entitlers.\*\*

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# METALLIZATION OF CARBON NANOTUBES FOR FIELD EMISSION APPLICATIONS

## TECHNICAL FIELD

The present invention relates in general to nanostructured materials, and in particular, to using modified carbon nanotubes for field emission applications.

#### BACKGROUND INFORMATION

Carbon annothes (CNTs) are currently being investigated for use as cold electors sources in a variety of applications. These include displays, microwave sources, x-ray tubes, etc. For CNTs to be used as a cold enthods, they must be piaced on a conductive surface (conductive substrates or conductive film on a nonconductive substrate). This has led some to phace catalysts on the substrate surface and grow the curbon annotables in stim using CVD techniques (Kim et al., st. Appl. Phys., 99(5), 2591 (2001)). However, this has several draw-backar. This techniques typically grows multi-wall earbon annotables (MWNTo). However, MWNTh have poorer finds emission quality compared to single-wall earbon annotables (WWNTo) (Kurachi et al., "FED with double-valled earbon annotable emitters," the 21st International Display Research Conference in Confunction with the 8<sup>th</sup> international Display Workshops, Nagoya Congress Center, Nagoya, Japan, Oct 16-19, 2001, pp. 1237-1240). The substrate is subjected to high temperature, typically above 600°C, limiting the ubstrates that can be used. Uniformity is difficult to achieve because of the high temperature growth processe required. As a result, the manufacence of esthodes using this process will be very expensive due to the number and complexity of post-processing steps needed to generate a material capable of producing the desired level of field emission.

Other investigations have content on processes for making CNT cathodes in a separate process, collecting them, and then dispensing them conto a substante using a variety of techniques (Kim et al., Disnond and Related Materials, 9, 1184 (2009)). This has several advantages over the in situ method described above. First, the fixtication of the CNT material is decoupled from the fixtication of the cathods. This permits choosing the optimal CNT material for the application (single-weal, double-well, multi-well, pourified, non-purified, etc.). Second, the dispensing processes is carried out a relatively low-temperatures, permitting greater fiscibility in the choice of substants. Third, uniform deposition over large area substantes in far more fiscible using currently-wealthleb, low-cost equipositic. Ournet dispensing processes, however, have their disadvantages. One of these is that the CNT fixers are often dispensed such that they clump together or are imbedded inside emotive material (Kim et al., "Toward a ridge of carbon namothe PEDs," the 21" international Display Research Conference in Conjunction with the 5" international Display Wachshops, Nagoya Congress Congres, Nagoya, Japan, Oct. 16-19, 2001, pp. 1221-1224). These factors limit the performance of the CNT material. "Activation" processes are often employed after dispensing the CNT material. These processes recover some of the reformance of the virgin CNT Change et al., U.S. Paters the, 6,454/22 IBI.). These "strivition" process steps, reformance of the virgin CNT Change et al., U.S. Paters the, 6,454/22 IBI.). These strivition "yourses steps,

however, can add cost to the product and may lead to non-uniform performance. Yet another disadvantage of current dispensing techniques is that the dispensed CNT fibers may not have sufficiently good contact to the subtrate or the substrate's conductive layer such that this impedes their ability to supply the electrons needed for field emission.

It has been recently found that by mixing CNT material with other nanoparticle materials, the field emission properties of the CNT were improved (Mao et al., U.S. Provisional Application No. 60417,246, incorporated herein by reference). Because neighboring manothers shidl the extracted electric fields from each other (Bosard et al., Adw. Man., 13, 184 (2001)), it is believed that this improvement is a result of induced separation of the CNT material by the nanoparticles. In situations where the CNT fibers are too close, they may electrically account the upplied electric field from each other. By increasing the separation between the fibers, the effective applied field strength at the emission situal is higher.

Many SWNT fibers are semiconducting with a bandgap that is dependent upon the chiral indices (a,m) of the SWNT. Choi et al. (US Patent 6,504,202 B.)) teach that, for field emission applications, this tendags be overcome by depositing a metal film on CNT fibers that are sineady stated to a substrate. Choi et al. teach that the CNT fibers are cooted after the fibers are grown using CVD techniques. This method has the inherent aforementioned disadvantages of growing CNTs on the substrate. Purthermore, were the CNT fibers to be dispensed onto the substrate and then coated, the problems of separating the CNT fibers for improved emission would still remain.

A method of aligning CNTs is disclosed in U.S. Patent No. 6,312,303 B1 to Yamiv et al. (incorporated herein by reference), whereby CNTs are aligned by including the CNTs in a host material, aligning the host material (such as liquid crystal material) and the host phase material then aligns the CNTs.

#### BRIEF DESCRIPTION OF THE DRAWINGS

For a more complete understanding of the present invention, and the advantages thereof, reference is now made to the following descriptions taken in conjunction with the accompanying drawings, in which:

- FIGURE 1 illustrates metallized carbon nanotubes on indium-tin-oxide (ITO)/glass, wherein the metal coating is not necessarily uniform over all of the carbon nanotubes (CNTs);
- FIGURE 2 illustrates an embodiment wherein metallized carbon nanotubes are magnetically-aligned while being dispensed;
  - FIGURE 3 illustrates a field emission display device incorporating the present invention;
  - FIGURE 4 illustrates an electroless plating bath used to coat carbon nanotubes with metal;
- FIGURE 5 illustrates field emission current vs. electric field for cobalt-coated and non-coated carbon pappings:

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FIGURE 6 illustrates an embodiment wherein a cathode substrate was placed on a set of six permanent magnets prior to dispensing magnetically-aligned metallized CNTs onto the substrate;

FIGURE 7 illustrates the arrangement of the six permanent magnets in FIGURE 6 prior to placing the ITO/glass substrate and dispensing the magnetically-aligned CNTs onto the substrate wherein the face of each of the block magnets is magnetized North-South as shown in the edge (side) view; and

FIGURE 8 illustrates field emission in a display device wherein the cathode comprises magneticallyactive metallized CNTs which were dispensed onto a substrate with magnets behind it, as in FIGURE 7.

#### DETAILED DESCRIPTION

The present invention is directed towards metallized carbon nanotubes, methods for making metallized carbon nanotubes; methods for dispensing metallized carbon nanotubes onto a substrate; methods for aligning metallized carbon nanotubes; cold enthode field emitting materials comprising metallized carbon nanotubes, aligned metallized carbon nanotubes, and combinations thereof; and methods of using metallized carbon nanotubes as cold cathods field emitters.

Metallized earbon manostubes, according to the present invention, are carbon nanotubes which have been at least partially conted with one or more metals. Carbon nanotubes, according to the present invention, include, but are not limited to, single-wall carbon ansotubes, multi-wall carbon annotubes, double-wall carbon manotubes, busclytubes, carbon fibrilis, derivatized carbon annotubes, chemically-modified carbon manotubes, metallic carbon nanotubes, metallic carbon nanotubes, metallic carbon nanotubes, and combinations threatof. Purily of the carbon annotube rescutar materials (i.e., the carbon nanotubes prior to being metallized) ranges generally from at least about 10 percent to at most about 100 percent, and more perificially from at least about 10 percent, and more perificially from at least should 20 percent to a ment about 100 percent, and more perificially from at least should 20 percent to a ment about 100 percent. The perificially from at least should 20 percent to a ment about 100 percent. A perificially from at least should 20 percent to a ment about 100 percent. The perificially from at least should 20 percent to a ment about 100 percent. The perificially from at least should 20 percent to a ment about 100 percent. The perificially from at least should 20 percent to a ment about 100 percent. The perificially from at least should 20 percent to a ment about 100 percent. The perificially from at least should 20 percent to a ment about 100 percent. The perificially from at least should 20 percent and a ment about 100 percent. The perificially from at least should 20 percent and a ment about 100 percent. The perificially from at least should 20 percent and a ment about 100 percent. The perificially from at least should 20 percent and a ment about 100 percent. The perificially from at least should 20 percent and a ment about 100 percent. The perificially from at least should 20 percent and a ment about 100 percent. The perificially from at least should 20 percent and a ment and a ment about 100 percent p

Motal costings (also termed "filma") on the carbon annotubes comprise one or more metal layers and agreemently in thickness from at least about 0.1 manometer (am), not at most about 10 microneters (am), specifically from at least about 0.1 manometer to at most about 1 microneters, and more specifically from at least about 0.5 manometers to at most about 1 microneters. Metal coatings on the carbon nanotubes include, but are not limited to nickel (NN), incn (Po), copper (Co), dilver (An), size (Zn), rhoddum (Rh), inc (Sn), cadmium (Co), beryllium (Bo), planlatum (Po), ballatum (Po), planlatum (Po), pl

metallized carbon annotable product ranges generally from at least about 0.1 percent to at most about 39 percent, specifically from at least about 1 percent to at most about 39 percent, and more specifically from at least about 5 percent to at most about 59 percent for a percent for a most about 59 percent for a most about 50 percent for a most form annotables for a most form for form for a most form for a most form for form for a most form for a most form annotable with any combination (s) of the aforement of most form for annotable s.

Exemplary methods of making mentilized outbon manotubes comprise the steps of a) providing a plurality of carbon nanotubes; b) preparing an electroless metal plating solution to form a reaction solution; d) adding said carbon nanotubes to said electroless metal plating solution to form a reaction solution; d) subjecting said reaction solution to a reducing condition which causes metal ions in solution to be reduced to metal and nucleats on the outbon nanotubes to produce metallized carbon annotubes; and c) removing said metallized carbon nanotubes from the reaction solution. In some embodiments of the present invention, the metallized carbon nanotubes are washed and dried after boing removed from the reaction solution.

Carbon nanotubes, as described herein, can be carbon nanotubes of any dimension, chirality, and unumber of walls that suitably provides for carbon nanotubes of the present invention and include, but are not limited to, single-wall carbon nanotubes (SWNTs), miltired to, single-wall carbon nanotubes (SWNTs), miltired to, single-wall carbon nanotubes (DWCTs), buckytubes, carbon fibrils, derivotized carbon nanotubes, chemically-modified carbon nanotubes, metallic carbon nanotubes, semiconducting carbon nanotubes, and combinations thereof. In some embodiments of the present invention, the carbon nanotubes are treated with hydrochloric acid prior to the metallization size.

An electroless plating solution (commonly refured to as a plating bath), according to the present invention, comprises a solvent, a metal sait, and a raducing agent (See Rammey et al., Electroless Plating and Costing of Methad; Noveys, Park Ridge, NI (1972), incorporated herein by reference, for a detailed description of electroless plating techniques). In some embediments of the present invention, there is a premoter species which helps to dissolve the metal sait. In some embediments, there may be a balancing agent to control the PII. The solvent can be any solvent which mixingly provides for the solvation of the electroless plating solution components. An examplary solvent is water. The metal sait can be any metal sait that suitably provides for electroless metal plating according to the present invention and includes, but is not limited to, saits of the following: nickel, iron, copper, silver, zinc, thedium, the, cadmium, chromium, bryllium, palladium, indrium, platinum, gold, and combinations thereof. In some sembodiments, alloys of two or race metals are plated on the cathon nanotwhere with this process. The reducing agent can be any reducing agent that suitably provides for the reduction of the metal sait scording to the present invention and includes, but is not limited to Natig-Oy-Ho, Natig-At-IC, Natig-At-IC, Natig-At-IC, and combinations thereof. The optional promoter

species can be any species which anisably promotes the electroless metal plating process of the present invention by facilitating the dissolution of the metal salt in the solution. Suitable promoter species include, but are not limited to C4f\_C/KN-4H\_C, Na\_C4f\_C/aN\_C4f\_C, Na\_C4f\_C, Na\_

In some embediments of the present invention, the process of adding the extroon anarotives to the electroses plating solution is carried out by first illuscuiteding the extron nanotives in a multiple solvent just prior to addition. This enhances their dispensal in the electroless plating solution to form a reaction solution. This reaction solution is subjected to a reducing condition which causes metal ions in solution to be reduced to metal and nucleate on the carbon manotives to produce metallized carbon nanotives. Reducing conditions, ecording to the present invention, are up conditions which suitably provide for a reduction of the metal loss in solution. Such reducing conditions, and upon the condition in solution. Such reducing conditions induce this reduction and include, but are not limited to, heating, irradiation, chemical activation, and combinations thereof. In some embodiments, the electroless plating solution is solution to the other on anothebra.

In some embediments of the present invention, the degree of carbon minoribe metalization (i.e., the amount of metal coated on the carbon nanotubes) is modulated by the amount of earbon nanotubes present in the reastion solution. In other embediments, the degree of carbon nanotube metaliation is modulated by the concentration of metal salts and reducing agents present in the reaction solution. In other embediments, the degree of carbon nanotube metaliation is modulated by the time the carbon nanotubes apend in the reaction solution. In still other embediments, a combination of one or more of the afterementioned methods of modulating the degree of carbon nanotube metaliation is used to produce a metalized carbon nanotube rounds with certain desirted characteristic dependent upon the degree to which it has been metalized.

In some embediments of the present invention, prior to the step of removing the metallized carbon manusobles from the reaction solution, a stabilizing agent is added to sinv the reduction of the metal ions. A stabilizing agent can be any species which satisfy provides for the slowing of the reduction process of the present invention and includes, but it not limited to, Grig-Oo, Crif-Oo, and combinations thereof. Such allowing of the reaction facilitates greater control over the names of the end product. Statistical methods of memoring the metallized carbon manufactor that the step of removing the metallized carbon manufactor thereof. In some embediments of the present invention, after the step of removing the metallized ordron manufactor of most fractions that the step of removing the metallized carbon manufactor of the metallized carbon manufactor product from the final product. Suitable solvents include, but are not limited to, water, injurgery) also combinations thereof. Optional driving of the metallized carbon manufactor product can be carried out by any drying process which satisfiely provides for the drying of the metallized carbon manufactor product can be carried out by any drying process which satisfiely provides for the drying of the metallized carbon manufactor according to the present invention and includes, but is not limited to, heating, exposure to vaccoun, revenue heating, including, and combinations thereof.

Exemplary methods of dispensing metallized carbon annotubes onto a substrate comprises: a) dispensing the metallized carbon annotubes in a solvent to form a suspension; and by applying the suspension to a substrate using an "applicator means." Solvents into which the metallized carbon annotubes are dispersed include, but are not limited to, inpropassol, methanol, acetone, water, ethanol, and combinations thereof. Methods of dispersing the metallized carbon annotabes in the solvent include, but are not limited to, strings, shaking, ultrasonic assistance, and combinations thereof. FYGURE 1 illustrates one embodiment of metallized carbon nanotubes 100 on a substrate 103.

An applicator means, according to the present invention, can be any method which suitably dispenses the suspension of metallized carbon nanotubes onto a substrate in a controlled manner. Such application can be uniform or non-uniform, and can vary considerably in terms of the thickness of the resulting film, or layer, of metallized carbon nanotubes on the substrate. Suitable applicator means include, but are not limited to, printing, dispensing, painting, spaying, brushing, and combinations thereof. Suitable printing methods include, but are not limited to, inkjet printing, screen printing, off-set printing, and combinations thereof. An exemplary applicator means comprises a spraying technique whereby the suspension of metallized carbon nanotubes is sprayed onto a surface using a sprayer. While not intending to be bound by theory, a sprayer, according to the present invention, can be a pump sprayer which rapidly pushes the suspension through a small orifice and, upon exiting said orifice, the suspension becomes an aerosol of small suspension droplets which are directed toward the substrate surface. Optionally, the substrate can be heated during the application process to prevent the running of excess solvent. Typically, the substrate, after having applied the metallized carbon nanotubes to its surface, is dried to remove any excess solvent. A substrate, as described herein, can be any substrate which suitably provides for a surface on which to dispense metallized carbon nanotubes according to the present invention and includes, but is not limited to, metals, ceramics, glass, semiconductors, coated surfaces, layered materials, and combinations thereof.

In some embodiments of the present investion, the metallized carbon annotables are dispensed onto a substrate while under the influence of a magnetically-selve coating, the metallized earbon annotables have a magnetically-selve coating, the metallized earbon annotables are magnetically-selve coating, the metallized earbon annotables are magnetically-selved coating, the metallized earbon annotables are magnetically-selved assequent to their being dispensed on a substrate. Magnetic alignment, according to the present invention, can be ableved with one or more magnets selected from the group consisting of permanent magnets, electromagnets, and combinations thereof. Various embodiments of the present invention comprise magnetic fields which include, but are not limited to, magnetic fields which include, but are not limited to, magnetic fields which set uniform, non-uniform, directed, multi-directional, introprise, anisotropic, continuous, pulsed, and combinations thereof. In some embodiments, a magnetic field is applied to an entire substrate while a dispensing lead is natered over the substrate surface. In some embodiments of the present invention, the substrate is amagnetic. Magnetic alignment, according to the present invention, in a very clean process in text, unlike existing alignment processes (Chang et al., U.S. Patent No. 6436,221), nothing is required to come into contact with the nanotube surface in order to generate such alignment. Purcharrores, in

some embodiments of the present invention, the magnetic alignment process can be "patterned" such that some regions of the nanothole layer dispensed on a substrate are aligned in one direction, and are dispect in other directions in other regions. These directions can be with magnetic "north" either vertical to the plane (up or down) or in the plane, or any combination of in-plane and out-of-plane (slanted). The alignment process can be earried out either during the earbon nanothole deposition or after deposition.

FIGURE 2 illustrates an embodiment wherein metallized carbon nanombes are magnetically-stiligent while being dispensed. Restring to FIGURS 2, a dispensing head 202, which dispenses magnetic, metallized CNTs and which moves in X under Y directions, is restored over substrate 2011. As the magnetic, metallized CNTs are forced through nozzie 203, they are aligned with a magnetic field generated by coils 204 and power supply 205. Dispensed drops 206 of magnetically-aligned metallized CNTs can thus deposited in any surangement or orientation on a substrate surface. Optionally, an additional magnet 209 comprising a magnetic core 208 can be used to further direct the dispensing process and orient the magnetic, metallized CNTs. In some embodiments, an optional conting 207 is applied to the substrate before commencing with dispensing of the magnetic, metallized CNTs.

In some embodimente, as described later, permanent magnets can be arranged on one or both sides of substante 201 to easier in aligning magnetically-entire metallized CNTs during dispensing. Magnetic fields can also be supplied by larger electromagnetic colls that do not move with the dispensing bead.

In some embodiments of the present invention, an electro-magnetic head is rastered over a surface, after magnetically-active metallized CNTs have been dispensed onto said substrate. Such rastering produces patterned alignment. In these embodiments, the electro-magnetic bead writes a pattern into a surface much like a read/write head writes a pattern to a magnetic surface of a disk in a data storage "hard drive" of a commutat.

In some embodiments of the present invention, metallic carbon manothes are dispensed with manoparticles in a manner described periodicyl for the dispensing of unusualized carbon nanothese with nanoparticles (United States Provisional Patest Application, Sorial No. 60417,246, incorporated herein by reference). Such nanoparticles have compositions which include, but are not limited to, metals, semirontals, fullerence, semiconductors, delectrics, cenancies, metalloids, glasses, polymers, and combinations thereof. In some embodiments, the manoparticles are magnetically sortive nanoparticles. In such embodiments, magnetically-active metallized carbon manothes are dispensed with magnetically-active nanoparticles. In such embodiments, local magnetic field strongth can be increased during the alignment process, thus leading to potentially higher degrees of alignment of the metallized carbon nanothes.

In some embodiments of the present invention, the metallized earbon nanotubes are used for fieldemission application. In some embodiments, these metallized earbon nanotubes are more suitable for fieldemission applications than carbon nanotubes without a metal costing. While not intending to be bound by theory, it is likely that when incorporated into a device for field emission applications, the metallized earbon nanotubes are better separated from one another, creating a carbon munotube arrangement of lower density that reduces the shielding effects contributed by neighboring carbon manolubes. Purthermore, said metal coatings likely educate the few of electrons in semiconducing carbon manolubes and at the nanolube-substrate junction. In some embodiments of the present invention involving field emission applications, the metallized carbon manotubes are dispensed onto a substrate using an application means, and the resulting substrate (with the metallized carbon manolubes) is used as the cathode in, for example, a field emission display. Other field emission applications in which metallized carbon manotubes can be used include, but are not limited to X-ray sources, electron sources, if armys, microwave tubes, and combinations thereof.

In some embediments of the present invention wherein metallized CNTs are disposued onto a substrate surface for use as a cathode in field emission application, an optional taping process can be used to "sectivate" the CNT layer and produce better field emission. In such embodiments, an atthenive film or tape is placed on top of the CNT layer such that the adhesive is put in contact with the CNTs. The tape is then removed at an appropriate angle such that the CNTs on the surface of the layer can be vertically aligned to further enhance field emission exproperties. Such activation has been described previously for field emission cathodes compristing non-metallized CNTs (Chang et al., U.S. Patent No. 6,436,221 B); Yaniv et al., U.S. Provisional Patent application Serial No. 60948,836; both of which are incorporated herein by reference). Embodiments using magnetic alignment may obvise this tens.

Thus, as disclosed herein, the present invention is also directed towards an improved field emission cathode using carbon nanotube emitters that are first coated with a metal film and then dispensed onto the cathode. This field emission cathode is illustrated in FIGURE 1. Referring to FIGURE 1, metallized carbon nanotubes 106 are shown on a substrate 103 which comprises a conductive layer 102 and an optional layer 101, which can be either conductive or non-conductive. Collectively, this forms field emission cathode 100. This cathode has advantages over the current art in that: a) the metal layer provides a high level of electrical conductivity along the length of the CNT fiber even if the fiber is semiconducting; b) the metal layer provides an additional means of separating the CNT fibers from each other, decreasing the mutual electrical shielding and eliminating the need for post-deposition activation steps; c) metal-coated earbon fibers adhere to metal layers on the substrate much more strongly than do bare carbon nanotubes (adhesion forces between metals are much stronger than the adhesion forces between the substrate and the un-metallized carbon nanotubes); and the metal coatings can be applied to SWNTs and MWNTs, semiconducting or metallic CNTs, purified or non-purified CNTs-all using standard electrolytic techniques permitting selection from a large variety of available CNT fibers. Furthermore, in some embodiments of the present invention, the improved field emission cathode comprises metallized CNTs which can be magnetically-aligned. Magnetic alignment of these metallized carbon nanotubes within the field emission cathode can be in any desired orientation, and can include any or all of the metallized carbon nanotubes. Alignment can be patterned or uniform. Improved field emission from nonmetallized carbon nanotube-based field emission cathodes has been realized when the nanotubes are vertically aligned (See United States Provisional Patent Application, Serial No. 60/348,856, incorporated herein by reference).

Referring to FIGURB 3, the field emission cathode described shows can be incorporated into field maken display 300. On substrate 301, conductive layer 302 is deposited and metallized exteron anonthole layer 303 is deposited on top thereof. The smool includes substrate 304, which may be a glass substrate, conductive layer 305, which may be indism-tin-existe, and a phosphor layer 306 for receiving electrons emitted from metallized ourbon amonous layer 303. Electrons are emitted from layer 303 in response to an appropriate electric field between the succelor afth the existion.

In some embediments of the present invention, earbon monthless are coated with a magneticallyactive, but non-mentilic species. Costed monobles such as these can be made by first depositing a metal
containg, as described above, and then resulting this costing with other chemicals, such as containt (e.g.,
covygen.), to form compounds that are no longer metallio, but which are still magnetic. In other embodiments,
such non-metallic magnetically-active coatings are chemically precipitated out of a solution cate the carbon
nanothless. An example of a non-metallic magnetically-active material which can be applied to carbon
nanothless as coating is magnetite (Fey.O). Methods of depositing magnetite in this manner are known in the
ref (Berger et al., "Preparation and Properties of an Aqueous Ferrofinial," J. Chem. Edu., 76(7), 943 (1999);
Palacin et al., "Patterning with Magnetic Materials at the Micros Scale," Chem. Mater., 8, 1316 (1990); both
of which are incorporated herein by reference). One suitable method of depositing magnetic on extro
menculous involves preparing an aqueous solution complicing a mixture of Felly and Fe(III). balks and then
reacting this with ammonium hydroxide in the presence of carbon nanothbes. The iron then precipitates out of
solution as Fe<sub>0</sub>O<sub>0</sub>, coating the carbon nanothbes in the process. A surfactant may be employed to ficilitate
dispersion of the outron nanothbes within this solution.

In other embodiments of the present investion, other types of annostructured materials can be used in place of earbon annotables. These other nanostructured materials can be metallized, dispensed on a substante, and, I metallized with a magnetically-sclor most, deep can be aligned—all in the same material which the metallized carbon nanotabes are. Such associates materials include, but are not limited to, boron zitide nanostructures materials include, but are not limited to, boron zitide nanostructures, and nanowires of silicon, silicon carbide, gallium mitride, indium phosphide, and combinations thereof.

The process, according to the present invention, of first conting CNTs with metal and than disposable them onto a substrate has a number of advantages, puriouslarly for filled emission applications. Such a method of disposable CNTs not a substrate syrers to inhibit disumping, provides for sufficiently good contact to the substrate, overcomes the limitations imposed by semiconducting CNTs, and it obviates the need for activation processes. Yet another advantage, in enrodiments where the metal conting is megnetically active, is the ability to slight the metal contact CNTs before, during, or after deposition.

The following examples are provided to more fully illustrate some of the embodiments of the present invention. The examples illustrate methods by which metal-countd (metallized) CNTs can be made and prepared for field emission applications. It should be appreciated by those of skill in the art that the techniques disclosed in the examples which follow represent techniques discovered by the six wentor to function well in the practice of the invention, and thus can be considered to constitute exemplary modes for its practice. However, those of skill in the art should, in light of the present disclosure, appreciate that many changes can be made in the specific embodiments which are disclosed and still obtain a like or similar result without departing from the spirit and scope of the invention.

#### EXAMPLES

## EXAMPLE 1. Coating single-wall carbon nanotubes with a Cobalt thin film

This process provides a way of depositing a metal thin film or coating on the surface of carbon nanotubes using an electroless plating technique. Using this relatively inexpensive and simple process, metallized carbon nanotubes can be made efficiently in relatively large amounts.

The single-wall earbon nanotube (SWNT) material used here was purchased from Iljin Nanotech, Inc. (Korea). The length of the SWNTs ranged from approximately several micrometers to approximately 20 micrometers, and the diameters were generally less than about 2 nanometers.

Referring to FIGURB 4, electroless plating apparatus 400 comprises an electroless plating solution 404 contained in a beater 403 which in turn is immersed in a wester total 402. Water both 402 is beated by a magnatio stirring hobjists 401 and temperature is monitored by thermometer 406. Stirring is accomplished with stir bear 405 activated by the magnatio stirring hobjists 401 and the stirring motor 407. In the present example, electroless plating solution 404 comprises water and the following chamicals:

- A cobalt (Co) salt (CoSO<sub>4</sub>·7H<sub>2</sub>O) to provide Co ions (Note that other salts may be used, e.g., CoCl<sub>2</sub>·6H<sub>2</sub>O). Concentration of this component is approximately 20-28 grams per liter.
- A reducing agent (NaH<sub>2</sub>PO<sub>2</sub>·H<sub>2</sub>O) to reduce Co ions to Co(0). Concentration of this component is approximately 18-25 grams per liter.
- A promoter species to facilitate dissolution of the Co salt into the solution (C<sub>4</sub>H<sub>6</sub>O<sub>6</sub>KNa·4H<sub>2</sub>O).
   Concentration of this component is approximately 140-160 grams per liter.
- A stabilizing agent (H<sub>3</sub>BO<sub>3</sub>), to slow the reducing reaction. Concentration of this component is 27-35
  grams per liter.
- A balancing agent (NaOH). This is used to control the pH value of the solution. The amount of this material that is used is that needed to maintain a pH of 8-10 for the metal plating solution.

The above chemicals were dissolved in deionized water up to 900 milliliters.

The cobait ions in this solution undergo reduction under a reducing condition of approximately \$5-95°C. The pH of the solution needs to be controlled before and during the reaction. In this example, the pH when was maintained at about 9. MoSH was added during the shaltne process to control the pH of the solution.

Approximately 3-4 grams of carbon nanotube powder is ultrasonicated in a beaker containing approximately 100 milliliters of water for several minutes before being introduced into the electroless plating

solution (after addition, total solution is 1000 millilliers). After the solution is prepared, it is bested in a water bath to 83-95°C and the ultrasonicated SWNTE are then added to the electroless plating solution quickly while the solution was strired. Because the carbon annotable easily clamp together, the water-CNT mixture should be ultrasonicated immediately before adding it to the plating solution. The typical reaction time in the plating solution is shout 5-10 minutes. Longer times do not appear to affect the results greatly. During the reaction, gas is evolved from the solution. The solution is pink at the beginning but gradually turns coloriess. At the end of the reaction, little or no gas is evolved from the solution.

After reaction/deposition of metal, the reaction beaker is taken out of the water that and allowed to cold own to rount temperature. After several minutes, die metallized carbon nanotube powders collect at the bottom of the beater and the solution is decanted from the powder. The powder is washed several time, each time being careful to not disturb the powder. Washing dilutes the concentration of any electrodes plating reactants still remaining on the powder after the reaction. The powder is then removed and deted in a furnment of the powder is then removed and deted in a furnment of the powder is then removed and deted in a furnment of the powder is then removed and deted in a furnment of the powder is then removed and deted in formation of the powder is the removed and deted in formation of the powder is the removed and deted in formation of the powder is the removed and deted in a furnment of the powder is the removed and deted in formation of the powder is the powder in the powder in the powder in the powder is the powder in the powder in the powder in the powder in the powder is the powder in the pow

#### EXAMPLE 2. Dispensing carbon nanotubes onto a substrate

In this example, cobalt-metallized SWMT provder was mixed with isopropyl alcobal (IPA) to form a uniquenion. The suspension comprised approximately I gram of metallized SWMTs in 1000m1 IPA. Because the SWMTs chump together readily, ultrasonic agitation was used to disperse the manothes in the IPA before spraying the solution cuto cathods substates. The SWMTIPA suspension was uprayed onto conductive indium-tim-oxide (ITO)/glass substates with an area of 2x2cm<sup>2</sup>. In order to prevent the IPA from flowing uncontrollably, the substates was bested up to approximately 30-70°C on both the front side and back side during the spraying process. The substate was sprayed back and forth several to tens of times until the carbon nanothes covered on the entire surface. The thickness of the carbon nanothes layer was about 1-20µm. The film was then dried in sir.

#### 3. Field emission test of the samples

Substantes with metallized SWNT material contect on them were prepared as cathodes and tested for idel enablish properties as illustrated in FIGURES 1 and 3. Non-metallized SWNT contect substants were also prepared in an identical fishion by the spray process for comparison purposes. The cathodes were tested by mounting them with a phospher screen in a dode configuration with a spr of about 0.5mm. The test assumbly was placed in a vacuum chamber and pumped to 10<sup>-70</sup> terr. The electrical properties of the cathodes were then measured by applying a negative, pulsed voilage to the cathodes and holding the nancle at ground potential and measuring the current at the anode. A pulsed voilage was used to prevent damage to the phosphor screen at the pulse current to 10<sup>-10</sup> terr. The control of the current of the cathodes were not "nectivated," they were tested as they were deposited. It was found that the metallized CNT cathodes were very table and very uniform. The non-metallized cathodes typically were unstable during the turn-on process (sevent large events occurred.) From FIGURES 5 it can be seen that metallized CNTN y yield much

better field emission properties than the non-metallized SWNTs. Tests on the cathodes show threshold extraction fields of about ZV/µm and emission current of 30mA at 4V/µm for Co-coated CNT compared with extraction fields of 3.5V/µm and emission current of 30mA at 6.5V/µm for non-metallized CNTs.

All of the compositions and methods disclosed and claimed herein can be made and executed without under experimentation in light of the present disclosure. While the compositions and methods of this invention have been described in terms of preferred embodiments, it will be apparent to those of feld lin the art that variations may be applied to the compositions and methods and in the steps or in the sequence of steps of the methods described herein without departing from the ocnept, spirit, and scope of the invention. More predictably, it will be apparent that contain segats which are both chemically and physiologically related may be substituted for the agents described herein while the same or similar results would be achieved. All such similar substitutes and modifications apparent to those delibed in the art are deemed to be within the spirit, scope, and concern of the invention as defined by the appended claims.

# 4. Dispensing and alignment of magnetically-active metallized carbon nanotubes

Cobalt-coated SWNTs were made according to the technique outlined in EXAMFLE 1. Referring to FIGURS 6, a 25cm x 25cm enabode substrate 602 comprising FTO/glass was placed on top of six signoent 10cm x 15cm permanent magnets 601 as shown in FIGURS 6. The magnetically-active metallized SWNTs were then spayed into the outhods substrate to form a CNT layer 603 in accordance with the technique outlined in EXAMPLE 2.

In some embodiments, a masking layer can be placed on the surface of the substrate to pattern the CNT layer during the spraying process. In some embodiments, this mask layer is a metal foil with holes that allow the CNTs to collect on the substrate is defined pattern. In some embodiments, the metal foil is also magnetic and is attracted to the magnets 601 on the other side of the substrate to hold the foil firmly to the cathods substrate 602. In some embodiments, the magnets foil can be represented in magnetic poles on the ends or adder or faces of the magnet. In this example, the poles were on the shees of the magnet as shown in PiGURB 7. The armagement of the magnets are also shown in FiGURB 7. Soll refarring to PiGURB 7, other armagements are possible, including a complete reversal of north to south and south to north. In some embodiments the substrate 602 itself is magnetically sective and in such embodiments magnets 601 may not be needed.

In the current example, after spanying the magnetically-active metallized SWNTs can the cathode substrate, the magnets are removed and the cathode incorporated into a field emission display device, as in FIGURE 3. FIGURE 8 illustrates this device in use. FIGURE 8 shows an image of a field emission display device which depicts field emission intensity (bright spots) on a phosphor arrora. It is interesting to note that the regions of highest intensity are those where there were adjoining magnets on the backside (See FIGURES 6 & 7).

## WHAT IS CLAIMED IS:

- A magnetically-active metallized carbon nanotube material comprising carbon nanotubes which have a
  magnetically-active metal coating on them.
- 2. The material of Claim 1, wherein the carbon manthbes are selected from the group consisting of single-wall carbon nanotubes, multi-wall carbon nanotubes, double-wall carbon nanotubes, buckyrubes, exbon librils, earthon nanotubes, metallic carbon nanotubes, metallic carbon nanotubes, metallic carbon nanotubes, metallic carbon nanotubes, as micromoducing carbon nanotubes, and combinations thereof.
- The material of Claim 1, wherein the metal coating has a thickness which ranges from at least about 0.1
  nanometers to at most about 1 micrometer.
- An apparatus comprising:
  - a) a substrate; and
  - b) magnetically-active metallized carbon nanotubes.
- The cathode of Claim 4, wherein at least some of the magnetically-active metallized carbon nanotubes are aligned,
- 6. The cathode of Claim 4, further comprising nanoparticles.
- 7. The cathode of Claim 6, wherein at least some of the nanoparticles are magnetically active.
- 8. A method comprising the steps of:
  - a) providing a substrate; and
  - dispensing magnetically-active metallized carbon nanotubes onto said substrate using an applicator means.
- The method of Claim 8, wherein the applicator means comprises a spraying technique whereby a suspension of magnetically-active metallized carbon nanohubes suspension in a suitable solvent is sprayed onto said substanto.
- 10. The method of Claim 8, wherein the magnetically-active metallized carbon nanotubes are dispensed onto said substrate with carbon nanotubes that are not magnetically active.
- 11. The method of Claim 8, wherein the magnetically-active metallized carbon nanotubes are dispensed onto said substrate with nanoparticles.

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- 2. The method of Claim 11, wherein at least some of the nanoparticles are magnetically active.
- 13. The method of Claim 8, wherein the magnetically-active metallized carbon nanotubes are aligned with a magnetic field while they are being dispensed.
- 14. The method of Claim 8, wherein the magnetically-active metallized carbon nanotubes are aligned with a magnetic field subsequent to being dispensed.
- The method of Claim 8, wherein the substrate is magnetically-active.
- A field emission display device comprising:
  - a) an anode assembly; and
  - b) a cathode assembly, wherein the cathode assembly comprises:
    - a substrate;
    - an electrically conducting layer deposited on the substrate; and
      - a layer of magnetically-active metallized carbon nanotubes deposited over the electrically conducting layer.
- 17. A metallized carbon nanotube material comprising carbon nanotubes which have a metal coating on them.
- 18. The material of Claim 17, wherein the carbon nanotubes are selected from the group consisting of single-wall carbon nanotubes, multi-wall carbon nanotubes, buckytubes, carbon fibrils, derivatized carbon nanotubes, chemically-modified carbon nanotubes, metallic carbon nanotubes, semiconducting carbon nanotubes, and combinations thereof.
- 19. The material of Claim 17, wherein the metal coating is uniformly distributed over an exterior surface of said carbon nanotubes.
- 20. The material of Chaim 17, wherein the motal is selected from the group consisting of nickel, iron, coopper, silver, zinc, rhodium, tin, cadmium, chromium, beryllium, palladium, indium, platnum, gold, and combinations thereof.
- 21. The method of Claim 17, wherein the metal coating has a thickness which ranges from at least about 0.1 nanometers to at most about 1 micrometer.
- A method of making metallized carbon nanotubes comprising the steps of:
  - a) providing a plurality of carbon nanotubes;
  - b) preparing an electroless metal plating solution;
  - adding said carbon nanotubes to said electroless plating solution;

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- d) subjecting said electroless plating solution to a reducing condition which causes metal ions in solution to be reduced to metal and nucleate on the carbon nanotubes to produce metallized carbon nanotubes: and
- e) removing said metallized carbon nanotubes from solution.
- 23. The method of Claim 22, further comprising the step washing the metallized carbon nanotubes.
- 24. The method of Claim 22, further comprising the step of drying the metallized carbon nanotubes.
- 25. The method of Chaim 22, wherein the carbon nanotubes are selected from the group consisting of single-wall carbon nanotubes, multi-wall carbon nanotubes, buckynizes, carbon fibrils, derivetized earbon nanotubes, chemically-modified carbon nanotubes, metallic earbon nanotubes, semiconducting earbon nanotubes, and combinations thereof.
- 26. The method of Claim 22, further comprising the step of treating the carbon nanotubes with hydrochloric acid prior to their metallization.
- 27. The method of Claim 22, wherein the electroless plating solution comprises a solvent, a metal sait, and a reducing agent.
- 28. The method of Claim 27, wherein the electroless plating solution further comprises an optional component selected from the group consisting of a promoter species, an inhibiting agent, a balancing agent, and combinations thereof.
- 29. The method of Claim 27, wherein the metal salt comprises a metal selected form the group consisting of nickel, iron, copper, silver, zino, rhodium, tin, cadmiture, chromium, beryllium, palladium, indium, piatnum, ped, and combinations thereof.
- 30. The method of Claim 22, wherein the step of adding said carbon nanotubes to said electroless plating solution further comprises ultrasonicating the carbon nanotubes in a solvent just prior to addition.
- The method of Claim 22, wherein the step of removing said metallized carbon nanotubes from solution further comprises a separation technique selected from the group consisting of filtration, centrifugation, and combinations thereof.
- Metallized carbon nanotubes made by a process comprising the steps of:
  - a) providing a plurality of carbon nanotubes;
  - b) preparing an electroless metal plating solution;
  - c) adding said carbon nanotubes to said electroless plating solution;
  - d) subjecting said electroless plating solution to a reducing condition which causes metal ions in solution to be reduced to metal and nucleate on the carbon nanotubes to produce metallized carbon nanotubes; and

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- e) removing said metallized carbon nanotubes from solution.
- A cathode for field emission applications comprising:
  - a) a substrate; and
  - b) metallized carbon nanotubes.
- A method of making cathodes for field emission applications comprising the steps of:
   a) providing a suitable substrate; and
  - b) dispensing metallized carbon nanotubes onto said substrate using an applicator means.
- 35. The method of Claim 34, wherein the applicator means comprises a spraying technique whereby a suspension of metallized carbon nanotubes suspended in a suitable solvent is sprayed onto said substrate.
- 36. The method of Claim 35, wherein the suspension of metallized carbon nanotubes is generated using ultrasonic assistance.
- A field emission display device comprising:
  - a) an anode which includes a phosphor deposited on a substrate; and
  - a cathode comprising a layer of metallized carbon on a substrate.
  - A field emission display device comprising:
    - a) an anode assembly; and
    - b) a cathode assembly, wherein the cathode assembly comprises:
      - a substrate;
      - an electically conducting layer deposited on the substrate; and
    - a layer of metallized carbon nanotubes deposited over the electrically conducting layer,
- The field emission display device of Claim 38, wherein the metallized carbon nanotubes comprise single-well carbon nanotubes.

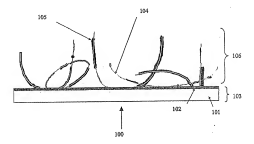


FIG. 1

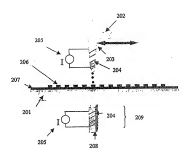


FIG. 2

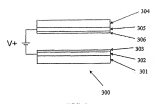


FIG. 3

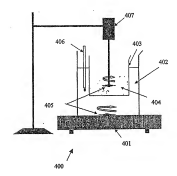


FIG. 4

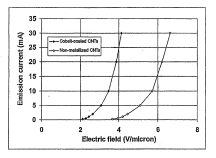


FIG. 5

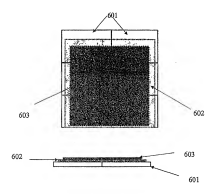


FIG. 6

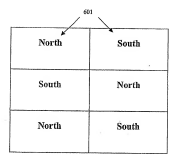




FIG. 7

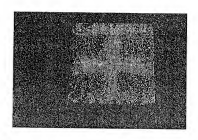


FIG. 8